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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.	
10/599,660	05/30/2007	Hoon Sunwoo	55326.15	2411	
	7590 03/02/200 O C/O BENNETT JON	EXAMINER			
1000 ATCO CENTRE			WEN, SHARON X		
10035 - 105 STREET EDMONTON, ALBERTA, AB T5J3T2		$\Gamma 2$	ART UNIT	PAPER NUMBER	
CANADA	·		1644		
			MAIL DATE	DELIVERY MODE	
			03/02/2009	PAPER	

Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

	Application No.	Applicant(s)			
Office Action Commence	10/599,660	SUNWOO ET AL.			
Office Action Summary	Examiner	Art Unit			
	SHARON WEN	1644			
The MAILING DATE of this communication ap Period for Reply	pears on the cover sheet with the	correspondence address			
A SHORTENED STATUTORY PERIOD FOR REPL WHICHEVER IS LONGER, FROM THE MAILING IDENTED TO THE	DATE OF THIS COMMUNICATIO .136(a). In no event, however, may a reply be tid d will apply and will expire SIX (6) MONTHS from te, cause the application to become ABANDONE	N. mely filed n the mailing date of this communication. ED (35 U.S.C. § 133).			
Status					
Responsive to communication(s) filed on 27 I This action is FINAL . 2b) ☐ This Since this application is in condition for allowed closed in accordance with the practice under	is action is non-final. ance except for formal matters, pr				
Disposition of Claims					
4) ☐ Claim(s) 1-9 and 14 is/are pending in the app 4a) Of the above claim(s) is/are withdra 5) ☐ Claim(s) is/are allowed. 6) ☐ Claim(s) 1-9 and 14 is/are rejected. 7) ☐ Claim(s) is/are objected to. 8) ☐ Claim(s) are subject to restriction and/	awn from consideration.				
Application Papers					
9) The specification is objected to by the Examin 10) The drawing(s) filed on is/are: a) ac Applicant may not request that any objection to the Replacement drawing sheet(s) including the correct 11) The oath or declaration is objected to by the E	cepted or b) objected to by the drawing(s) be held in abeyance. Section is required if the drawing(s) is ob	ee 37 CFR 1.85(a). ojected to. See 37 CFR 1.121(d).			
Priority under 35 U.S.C. § 119					
 12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f). a) All b) Some * c) None of: 1. Certified copies of the priority documents have been received. 2. Certified copies of the priority documents have been received in Application No. 3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)). * See the attached detailed Office action for a list of the certified copies not received. 					
Attachment(s) 1) Notice of References Cited (PTO-892) 2) Notice of Draftsperson's Patent Drawing Review (PTO-948) 3) Information Disclosure Statement(s) (PTO/SB/08) Paper No(s)/Mail Date	4) Interview Summary Paper No(s)/Mail D 5) Notice of Informal I 6) Other:	oate			

Application/Control Number: 10/599,660 Page 2

Art Unit: 1644

DETAILED ACTION

1. Applicant's amendment, filed 11/27/2008, has been entered.

Claims 10-13 have been canceled.

Claim 14 has been added.

Claims 1-9 and 14 are pending and currently under examination as they read on a therapeutic composition comprising anti-gluten egg yolk antibodies wherein the specific anti-gluten antibody reads on the elected anti-gliadin antibody.

2. This Action will be in response to Applicant's Arguments/Remarks, filed 11/27/2008.

The rejections of record can be found in the previous Office Action, mailed 08/13/2008.

Ora Mune Magna

3. Applicant's statement on Ora Mune Magna product which indicated that it was provided to the supplier by Applicant *no earlier* than May, 2006 has been acknowledged.

Claim Rejections - 35 USC § 103

- 4. The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:
 - (a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.
- 5. Claims 1-9 and 14 are rejected under 35 U.S.C. 103 (a) as being unpatentable over Lee (U.S. Patent 5,367,054, reference of record) in view of Ellis et al. (*Gut* 1998, 43:190-195, reference of record).

Art Unit: 1644

Applicant's argument has been considered in full but has not been found convincing essentially for reasons of record and reiterated herein for Applicant's convenience.

The present claims are drawn to a composition comprising anti-gluten IgY antibodies. Ellis et al. teach anti-gluten antibodies that are raised against gliadin, the elected species of anti-gluten antibody, wherein the antibodies are polyclonal and monoclonal IgG antibodies (see entire document, in particular, see page 190, last paragraph on the right column).

The difference between the teaching of Ellis and the present claims is that Ellis's antibodies are not IgY antibodies. However, raising IgY antibodies was a well-known technology in the art at the time of the invention was made as evidenced by Lee (see entire document, in particular, see Background of the Invention). In particular, Lee teaches the process of producing egg yolk antibodies by 1) immunizing the egg-laying fowl with the antigen of interest; 2) collect eggs from the immunized fowl; and 3) prepare the composition from the egg yolk or IgY purified from the egg yolk (see e.g., column 8, lines 7-15, and Figure 1). Moreover, Lee teaches the egg yolk is liquid or dried in the purification process (see Figure 1 and column 3, lines 51-57).

Given that Ellis et al. teach using the anti-gliadin antibody containing composition for immunodetection and the teaching by Lee on the advantage of making IgY antibodies i.e., that egg yolk is a very good source of specific antibodies and that the antibodies are more specific (see column 1, lines 34-46), one of ordinary skill in the art would have been motivated to make the anti-gliadin antibodies for immunodetection as IgY antibodies for the high specificity offered by the egg yolk.

Moreover, ordinary skill in the art would have reasonable expectation of success in making the anti-gliadin IgY antibodies in view of the detailed procedures in making and purifying IgY antibodies outlines in Lee (see e.g., column 8, lines 7-15, and Figure 1) and the detailed disclosure of how to prepared the gliadin antigen taught by Ellis et al. (see page 191, left column).

In view of the composition comprising the anti-gluten antibodies taught by Ellis et al. (see paragraph bridging pages 191-192) and the advantage and practicality in IgY production taught by Lee (see Background of the Invention and Figure 1), it would have been *prima facie* obviate to make an anti-gluten IgY antibody.

Regarding the product-by-process limitation for producing the antibody provided by the present claims, is noted that such process does not distinguish from the antibody in the art.

"[E]ven though product-by-process claims are limited by and defined by the process, determination of patentability is based on the product itself. The patentability of a product does not depend on its method of production. If the product in the product-by-process claim is the same as or obvious from a product of the prior art, the claim is unpatentable even though the prior product was made by a different process." *In re Thorpe*, 777 F.2d 695, 698, 227 USPQ 964, 966 (Fed. Cir. 1985).

Art Unit: 1644

Furthermore, it is noted that the claims provide intended uses for the composition comprising the antibody (i.e., for treating celiac disease and for oral administration) but such intended uses do not distinguish from the composition in the art. See e.g. MPEP § 2114.

In response to Applicant's argument that the combination of Ellis and Lee do not render obvious the claimed composition comprising and anti-gluten IgY antibody because Ellis and Lee do not teach the particular therapeutic use, i.e., for treating celiac disease, it is noted that the intended use for the composition do <u>not</u> distinguish from the composition in the art.

The rationale to support a conclusion that the claims would have been obvious is that all the claimed elements (e.g., anti-gluten antibody / IgY antibodies) were known in the prior art and one skilled in the art could have arrived at the claimed invention by using known methods (making an anti-gluten antibody and making an IgY antibody) with no change in their respective functions and the combination would have yielded nothing more than predictable results of making an anti-gluten IgY antibody.

The rationale to support a conclusion that the claims would have been obvious is that a particular known technique (making IgY antibodies) was recognized as part of the ordinary capabilities of one skilled in the art. One of ordinary skill in the art would have been capable of applying this known technique to a known product (e.g. anti-gluten antibody) that was ready for improvement and the results would have been predictable to one of ordinary skill in the art.

Given that antibodies were well-known in the art for their pharmaceutical and therapeutic uses, it would have been obvious to one of ordinary skill in the art, at the time of the invention was made to make an composition comprising an anti-gluten IgY antibody for pharmaceutical or therapeutic purposes because IgY antibodies were known to offer advantages over the conventional antibodies as taught by Lee (see, e.g., column 1, lines 35-47). In particular, IgY antibodies are relatively easy to produce and have high specificity. Moreover, IgY antibodies provide oral routes of administration. Therefore, one of ordinary skill would have been motivated to make an anti-gluten IgY antibody given the advantages of IgY antibodies over the conventional antibodies.

Application/Control Number: 10/599,660 Page 5

Art Unit: 1644

Furthermore, it would have been obvious to one of ordinary skill in the art to include a physiologically acceptable carrier, excipient or diluent, which reads on water or any physiological buffer.

Applicant's arguments have not been persuasive.

Therefore, the rejection of record is **maintained** for the reasons of record, as it applies to the newly added and amended claims.

Conclusion

- 6. No claim is allowed.
- 7. **THIS ACTION IS MADE FINAL.** Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the mailing date of this final action.

8. Any inquiry concerning this communication or earlier communications from the examiner should be directed to SHARON WEN whose telephone number is (571)270-3064. The examiner can normally be reached on Monday-Thursday, 8:30AM-6:00PM, ALT. Friday, EST.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Eileen O'Hara can be reached on (571)272-0878. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

Application/Control Number: 10/599,660 Page 6

Art Unit: 1644

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/Sharon Wen/ Examiner, Art Unit 1644 February 24, 2009

/Phillip Gambel/
Primary Examiner
Technology Center 1600
Art Unit 1644
February 26, 2009